GenCore version 6.3 Copyright (c) 1993 - 2009 Biocceleration Ltd.

OM nucleic - nucleic search, using sw model

Run on: June 26, 2009, 03:22:05; Search time 25 Seconds (without alignments)

20013.649 Million cell

updates/sec

Title: US-09-300-482-298

Perfect score: 388 Sequence: 1

ggagaaagaaagaaagatg.....ttaaagcctgggacccccgt 388

Scoring table: IDENTITY_NUC

Gapop 10.0 , Gapext 1.0

Searched: 1681669 segs, 647077473 residues

Total number of hits satisfying chosen parameters: 3363338

Minimum DB seg length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Published_Applications_NA_New:*

Ι:

/ABSS/Data/CRF/ptodata/2/pubpna/US09_NEW_PUB.seq:*
2:

/ABSS/Data/CRF/ptodata/2/pubpna/US10_NEW_PUB.seq:*

/ABSS/Data/CRF/ptodata/2/pubpna/US11_NEW_PUB.seq:*
4:

/ABSS/Data/CRF/ptodata/2/pubpna/US12 NEW PUB.seg:*

 $\,$ Pred. No. is the number of results predicted by chance to have a

score greater than or equal to the score of the result being printed, $% \left(1\right) =\left(1\right) \left(1\right) +\left(1\right) \left(1\right) \left(1\right) +\left(1\right) \left(1\right) \left($

and is derived by analysis of the total score distribution.